

Section II (Amendments to the Claims)

Please amend claims 1, 11-14 and 17, and add new claims 19-21, as set out in the listing of claims 1-21 below.

1. (Currently Amended) A F_v antibody construct having binding sites for an CD16 receptor and a CD30 surface protein.
2. (Previously presented) The F_v antibody construct according to claim 1, wherein the CD16 receptor is derived from NK cells.
3. (Previously presented) The F_v antibody construct according to claim 1, wherein the CD30 surface protein is derived from a member selected from the group consisting of: Hodgkin's disease cells or Reed-Sternberg cells.
4. (Previously presented) The F_v antibody construct according to claim 1, wherein one binding site is present each.
5. (Previously presented) The F_v antibody construct according to claim 4, encoded by the expression vector pKTD16-30 (DEM 12960).
6. (Previously presented) The F_v antibody construct according to claim 1, wherein two binding sites are present for each.
7. (Previously presented) An expression vector, coding for the F_v antibody construct according to claim 1.
8. (Previously presented) The expression vector according to claim 7, which is pKID16-30 (DSM 12960).
9. (Previously presented) A transformant, containing the expression vector according to claim 7.

10. (Previously presented) A method of producing the F_v antibody construct according to claim 1, comprising culturing the transformant according to claim 9 under suitable conditions.

11. (Currently amended) A kit comprising:

(a) an F_v antibody construct according to the invention having binding sites for an CD16 receptor and a CD30 surface protein

and/or

(b) an expression vector according to the invention coding for said F_v antibody construct, and

(c) common at least one auxiliary substances, such as substance selected from the group consisting of buffers, solvents, carriers, controls and markers, wherein one or more representatives of the individual components may be present.

12. (Currently amended) Use of the F_v antibody construct according to claim 1 A method for lysis of cells expressing CD30 surface proteins, said method comprising contacting said cells with an F_v antibody construct having binding sites for an CD16 receptor and a CD30 surface protein.

13. (Currently amended) Use A method according to claim 12, wherein the cells are tumor cells.

14. (Currently amended) Use A method according to claim 13, wherein the tumor cells are selected from the group consisting of: Hodgkin's disease cells or Reed-Sternberg cells.

15. (Previously presented) The F_v antibody construct according to claim 2, wherein the CD30 surface protein is derived from a member selected from the group consisting of: Hodgkin's disease cells or Reed-Sternberg cells.

16. (Previously presented) An expression vector, coding for the F_v antibody construct according to claim 15.

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17. (Currently amended) Use of the F_v antibody construct according to claim 15 A method for lysis of cells expressing CD30 surface proteins, said method comprising contacting said cells with an F_v antibody construct having binding sites for an CD16 receptor and a CD30 surface protein, wherein the CD16 receptor is derived from NK cells, and wherein the CD30 surface protein is derived from a member selected from the group consisting of: Hodgkin's disease cells or Reed-Sternberg cells.

18. (Previously presented) A transformant, containing the expression vector according to claim 8.

19. (New) The F_v construct of claim 1, wherein said F_v antibody construct comprises elements (a) and (b) joined via a peptide linker:

- (a) a VH domain of an antibody and a VL domain of an anti-CD30 antibody, the domains being joined by a peptide linker; and
- (b) a VH domain of an anti-CD30 antibody and a VL domain of an anti-CD16 antibody, the domains joined by a peptide linker.

20. (New) A method of treatment of a tumor comprising the step of administering the F_v antibody construct according to claim 1.

21. (New) The method of claim 20, wherein the treatment comprises the lysis of Hodgkin's disease or Reed-Sternberg cells.